
Sequence Listing could not be accepted due to errors.

See attached Validation Report.

If you need help call the Patent Electronic Business Center at (866)

217-9197 (toll free).

Reviewer: Anne Corrigan

Timestamp: [year=2008; month=1; day=23; hr=13; min=45; sec=43; ms=375;]

Reviewer Comments:

<120> A METHOD FOR CREATING A FUNCTIONALLY ACTIVE CHIMERIC TYPE IIG
RESTRICTION ENDONUCLEASE (as amended)

The first line of the above <120> response exceeds the Sequence Rules' required 72-character line (this includes white spaces). Please insert a hard return after "TYPE" on the first line, and align the second line.

(from Sequence 2)

Val Asp Glu Ala Leu Leu Ile Lys Tyr His Gly Phe Ser Glu Lys Glu
515 520 525

Val Lys Gln Leu Arg Gly Ile Trp Lys Lys Leu Ser Gln Arg Arg Asn 530 540

Asn Arg Thr Lys Lys 545

Please delete the excess blank lines above: only one blank line should separate each amino acid line.

<210> 38

<211> 103

<212> PRT

<213> artificial

<220>

<223> segment of protein sequence of catechol O-methyltransferase

<400> 39

Please change the above <210> response to "39." <210> 38 was already shown.

Validated By CRFValidator v 1.0.3

Application No: 10800946 Version No: 2.0

Input Set:

Output Set:

Started: 2008-01-09 14:09:09.990

Finished: 2008-01-09 14:09:12.054

Elapsed: 0 hr(s) 0 min(s) 2 sec(s) 64 ms

Total Warnings: 25

Total Errors: 2

No. of SeqIDs Defined: 43

Actual SeqID Count: 43

Error code		Error Descript	ion								
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W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(20)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(21)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(22)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(23)
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W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(25)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(26)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(27)
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W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(29)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(30)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(31)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(32)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(33)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(34)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(35)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(36)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(37)
W	213	Artificial	or	Unknown	found	in	<213>	in	SEQ	ID	(38)

Input Set:

Output Set:

Started: 2008-01-09 14:09:09.990 **Finished:** 2008-01-09 14:09:12.054

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Error code Error Description

This error has occured more than 20 times, will not be displayed

E 212 Invalid Sequence ID Number; Expected 39 as next SeqID but skipped

E 212 Invalid Sequence ID Number; Expected 39 as next SeqID but skipped

```
<110> Xu, Shuang-yong
      Kobbe, Daniela
      Zhu, Zhenyu
      Samuelson, James
<120> A METHOD FOR CREATING A FUNCTIONALLY ACTIVE CHIMERIC TYPE IIG RESTRICTION
                                                                                    ENDONUCLEASE
(as amended)
<130> NEB-183-CIP
<140> 10800946
<141> 2004-03-15
<150> 10/150,028
<151> 2002-05-17
<150> 09/693,146
<151> 2000-07-02
<160> 43
<170> PatentIn version 3.2 (1-26) and 3.4 (27-43)
<210> 1
<211> 1650
<212> DNA
<213> Bacillus pumilus
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<222> (1)..(1650)
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Met Asn Gln Leu Ile Glu Asn Val Asn Leu Gln Lys Leu Arg Gly Gly
tat tac acc cct aaa gtt att gct gac ttt tta tgt caa tgg agt att
                                                                  96
Tyr Tyr Thr Pro Lys Val Ile Ala Asp Phe Leu Cys Gln Trp Ser Ile
             20
                                 25
caa gat gac aca aag agt gta ctt gaa ccc agt tgt gga gat ggt aat
                                                                  144
Gln Asp Asp Thr Lys Ser Val Leu Glu Pro Ser Cys Gly Asp Gly Asn
         35
ttt att gaa tcg gca ata ctt agg ttc aaa gaa ctt agt ata gat aat
                                                                  192
Phe Ile Glu Ser Ala Ile Leu Arg Phe Lys Glu Leu Ser Ile Asp Asn
                         55
gaa caa ctt aaa gga aga att aca gga gta gag cta att gaa gaa gaa
Glu Gln Leu Lys Gly Arg Ile Thr Gly Val Glu Leu Ile Glu Glu Glu
65
                     70
                                         75
                                                             80
```

288

gct ttg aaa gtt caa aat cga gca aat gag ttg ggg gtt gat aaa aac

Ala Leu Lys Val Gln Asn Arg Ala Asn Glu Leu Gly Val Asp Lys Asn

90 95

		-	aat Asn 100	_	_					_		_		_		336
			gat Asp										_			384
			gaa Glu			-	-		-	_	-	_	_		-	432
			aaa Lys									-				480
, ,	-		gct Ala					_			_	_	-	_	_	528
	_	-	gaa Glu 180			_	Ī	_		-	_	_		_		576
			aag Lys													624
	-		gaa Glu				_	-	-					_	-	672
Lys 225	Lys	Val	aat Asn	Lys	Gly 230	Lys	Gly	Ile	Arg	Val 235	Ile	Glu	Cys	Glu	Asn 240	720
			tta Leu													768
_			att Ile 260	-		_		_	_			_				816
	_	_	gaa Glu						_		_	_	_		_	864
-			tgt Cys		-			_		-	-			-		912
	_		gaa Glu			_	_		-			-		-		960

305 310 315 320

		gaa Glu	-					_						_		1008
		ata Ile				Ĩ.		-			-			_	Ť.,	1056
	_	gca Ala 355					_			_	_	_		_	_	1104
	_	att Ile		-					_			-	-			1152
		caa Gln					_		_		_					1200
		aga Arg		-		Ī	-		-		_	_	-	-		1248
		aaa Lys														1296
		agg Arg 435	-	-			-				_	_		-	_	1344
		ttt Phe														1392
-	-	tat Tyr				-	_			-			-			1440
-		cta Leu					_					_		-		1488
	-	gcc Ala			-	-	_			-		-		-		1536
-	-	gaa Glu 515	-									-			-	1584
_		cag Gln		_				_				_	_	_		1632

530 535 540

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Lys Lys Val Asn Lys Gly Lys Gly Ile Arg Val Ile Glu Cys Glu Asn

230 235 225 240 Leu Asp Gly Leu Asn Ser Ile Asp Phe Val Ala Ile Asn Gly Ser Asn 250 245 Val Lys Pro Ile Glu His Arg Thr Glu Lys Trp Thr Lys Tyr Phe Leu 260 265 Asn Glu Asp Glu Ile Leu Leu Gln Ser Leu Lys Glu Asp Lys Arg 280 275 Val Lys Asn Cys Asn Asp Tyr Phe Lys Thr Glu Val Gly Leu Val Thr 295 Gly Arg Asn Glu Phe Phe Met Met Lys Glu Asn Gln Val Lys Glu Trp 310 315 Asn Leu Glu Glu Tyr Thr Ile Pro Val Thr Gly Arg Ser Asn Gln Leu 325 330 Lys Gly Ile Thr Phe Thr Glu Asn Asp Phe His Glu Asn Ser Met Glu 340 345 Gln Lys Ala Ile His Leu Phe Leu Pro Pro Asp Glu Asp Phe Glu Lys 355 360 Leu Pro Ile Glu Cys Gln Asn Tyr Ile Lys Tyr Gly Glu Glu Lys Gly 375 Phe His Gln Gly Tyr Lys Thr Arg Ile Arg Lys Arg Trp Tyr Ile Thr 385 390 395

Pro Ser Arg Trp Val Pro Asp Ala Phe Ala Leu Arg Gln Val Asp Gly 405 410 415

Tyr Pro Lys Leu Ile Leu Asn Glu Thr Asp Ala Ser Ser Thr Asp Thr 420 425 430

Ile His Arg Val Arg Phe Lys Glu Gly Ile Asn Glu Lys Leu Ala Val
435 440 445

Val Ser Phe Leu Asn Ser Leu Thr Phe Ala Ser Ser Glu Ile Thr Gly 450 455 460

Arg Ser Tyr Gly Gly Gly Val Met Thr Phe Glu Pro Thr Glu Ile Gly 465 470 475 480

Glu Ile Leu Ile Pro Ser Phe Asp Asn Leu Ser Ile Asp Phe Asp Lys
485 490 495

Ile Asp Ala Leu Ile Arg Glu Lys Glu Ile Glu Lys Val Leu Asp Ile
500 505 510

Val Asp Glu Ala Leu Leu Ile Lys Tyr His Gly Phe Ser Glu Lys Glu 515 520 525

Val Lys Gln Le 530 Asn Arg Thr Ly 545	-	Ile Trp Ly 535	-	Ser Gln Arg 540	Arg Asn									
<210> 3 <211> 3030 <212> DNA <213> Bacillus pumilus														
<220> <221> CDS <222> (1)(3030)														
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ttt tta aaa co Phe Leu Lys P:		Asn Glu Th												
gac cca ctt ct Asp Pro Leu Le 35														
aaa aca cat at Lys Thr His II	-	3	_	-	-									
aaa gat gag ga Lys Asp Glu G														
ggt acg aga aa Gly Thr Arg Ly	-			_										
ttg aaa tca go Leu Lys Ser Ai	la Lys Ala	_	n Thr Arg											
gct aac ctt go Ala Asn Leu G 115		_			-									
tat gat tgt ag Tyr Asp Cys A: 130	_	-	s Ser Asp		· ·									

aga tat aaa gtt ttc tct tac gag gaa tat gaa gaa gca ttt gat gaa 480

Arg 145	Tyr	Lys	Val	Phe	Ser 150	Tyr	Glu	Glu	Tyr	Glu 155	Glu	Ala	Phe	Asp	Glu 160	
	_	_					gag Glu		_				_	_	_	528
-	_		-	-			aga Arg	_		-	_		-			576
		_					tgg Trp 200	_	_	_		_			_	624
					_		ggt Gly	_		_	-				-	672
	_				_		att Ile			_	_	_	_	_	_	720
		-			-		att Ile		-						-	768
		-	_			_	tct Ser	_								816
	-			-	-	_	ctc Leu 280	-			-	-		-	-	864
	-	_		-			agt Ser	_						_		912
	-			-	_	_	cca Pro				_	_			_	960
_					-		att Ile		, ,							1008
	1				_	_	gcg Ala					_	_			1056
				-	-	-	ata Ile 360			-			_			1104
_	-						gag Glu		_					_	_	1152

370 375 380